EAST Search History

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
1.1		(bi\$2arsen\$ or (bil or bis! or dil) adj (arsen\$ or arsa)).clm.	US-PGPUB; USPAT	OR	ON	2006/06/29 10:13
L2	8	(tetra\$1cystein\$ or tetra adj cystein\$ or ccxxcc).clm.	US-PGPUB; USPAT	OR	ON	2006/06/29 10:13

Checked L1, L2 SEL 6-29-2006 Library Sequence Search Mortony

GenCore version 5.1.9 Copyright (c) 1993 - 2006 Biocceleration Ltd.

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:24:38; Search time 203 Seconds

(without alignments)

77.464 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2849598

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : UniProt 7.2:*

1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

D = = 3 to		8				
Result	_	Query	_			
No.	Score	Match	Length	DB	ID	Description
						
1	53	52.5	157	1	VE6_HPV12	P36803 human papil
2	51	50.5	243	2	Q46MX5 RALEJ	Q46mx5 ralstonia e
3	51	50.5	278	2	Q6IGH0_DROME	Q6igh0 drosophila
4	51	50.5	502	2	Q9BGM9_9MAMM	Q9bgm9 tachyglossu
5	51	50.5	1408	2	Q381X2_9TRYP	Q381x2 trypanosoma
6	50	49.5	589	2	Q2JB13_9ACTO	Q2jb13 frankia sp.
7	50	49.5	1370	1	ZMYM3_HUMAN	Q14202 homo sapien
8	50	49.5	1370	1	ZMYM3_MOUSE	Q9jlm4 mus musculu
9	49.5	49.0	602	2	Q75NZ5_CHLRE	Q75nz5 chlamydomon
10	49	48.5	115	1	ALK1_PIG	P22298 sus scrofa
11	49	48.5	155	2	Q9PXB1_HPV08	Q9pxb1 human papil
12	49	48.5	168	1	VE6_HPV21	P28832 human papil
13	49	48.5	1067	2	Q4QFE4_LEIMA	Q4qfe4 leishmania
14	48.5	48.0	390	2	Q4S604_TETNG	Q4s604 tetraodon n
15	48	47.5	62	2	Q4PN38_IXOSC	Q4pn38 ixodes scap

```
P97430 mus musculu
Q548x8 mus musculu
Q548x8 mus musculu
Q40617 human papil
Q81986 human papil
Q913v6 human papil
Q64fq2 arabidopsis
Q62h93 burkholderi
Q4ivt4 azotobacter
Q9r0z8 rattus norv
P22422 human papil
P28830 human papil
P28830 human papil
Q8vmh1 pseudomonas
Q9byq5 homo sapien
Q9byr0 homo sapien
Q52396 pseudomonas
Q3e2m8 chloroflexu
Q3inu5 natronomona
Q341e3 rhodopseudo
Q4p5x7 ustilago ma
P16343 lonchocarpu
Q52509 pseudomonas
Q4ckp2 trypanosoma
Q37b06 rhodopseudo
Q618g9 homo sapien
Q8mz55 drosophila
P36806 human papil
Q9byr2 homo sapien
Q28583 ovis aries
Q3vgi5 sphingopyxi
Q2i0e2 oryza sativ
Q65t35 mannheimia
P91666 drosophila
P19922 bradyrhizob
Q4t2b4 tetraodon n
16
          48
                   47.5
                               131 1 ALK1 MOUSE
                                                                                  P97430 mus musculu
17
            48
                   47.5
                                131 2
                                            Q548X8 MOUSE
                                                                                 Q548x8 mus musculu
                              157 2
                                            O40617 HPVR7
18
            48
                   47.5
                   47.5 157 2
                                            Q81986 HPV05
19
            48
                                           Q913V6_9PAPI
Q64FQ2_ARATH
                   47.5 157 2
47.0 525 2
            48
20
         47.5
21
         47.5 47.0 676 2 O48785 ARATH
22
23
                   46.5
                               88 2 Q62H93 BURMA
          47
                              101 2 Q4IVT4_AZOVI
                   46.5
24
            47
                               131 2 Q9R0Z8 RAT
25
            47
                   46.5
           47 46.5 156 1 VE6_HPV47
47 46.5 171 1 VE6_HPV14
47 46.5 181 2 Q8VMH1_PSEPU
26
27
                                            Q8VMH1_PSEPU
28
            47 46.5 193 1
29
                                            KR415 HUMAN
30
            47
                   46.5 210 1
                                            KRA47 HUMAN
                                            Q52396 PSEST
31
            47
                   46.5 219 2
            47
                              330 2 Q3E2M8 CHLAU
32
                   46.5
           47 46.5 399 2 Q3INU5_NATPD
47 46.5 438 2 Q341E3_RHOPA
47 46.5 1175 2 Q4P5X7_USTMA
33
34
35
36
            46 45.5 80 1 IBB4 LONCA
37
            46 45.5
                               88 2 Q52509 PSESX
            46 45.5
                               95 2 Q4CKP2 TRYCR
38
            46 45.5 100 2 Q37B06_RHOPA
46 45.5 129 1 KRA56_HUMAN
39
40
                                            Q8MZ55_DROME
41
            46 45.5 161 2
42
            46 45.5 166 1 VE6 HPV19
43
            46 45.5 186 1
                                            KRA45 HUMAN
           46 45.5 191 2 Q28583_SHEEP
46 45.5 203 2 Q3VGI5_9SPHN
44
45
            46 45.5 232 2 Q2I0E2_ORYSA
46 45.5 298 2 Q65T35_MANSM
46
47
48
            46 45.5 412 2 P91666_DROME
            46 45.5 465 1 HYIN2 BRAJA
49
50
                   45.5 491 2 Q4T2B4 TETNG
            46
                                                                                  Q4t2b4 tetraodon n
```

```
VE6 HPV12
ID
     VE6 HPV12
                    STANDARD;
                                    PRT;
                                           157 AA.
AC
     P36803;
     01-JUN-1994, integrated into UniProtKB/Swiss-Prot.
DT
DT
     01-JUN-1994, sequence version 1.
DT
     07-FEB-2006, entry version 27.
DE
     Protein E6.
GN
     Name=E6;
OS
     Human papillomavirus type 12.
OC
     Viruses; dsDNA viruses, no RNA stage; Papillomaviridae;
OC
     Betapapillomavirus.
OX
     NCBI_TaxID=10604;
RN
RP
     NUCLEOTIDE SEQUENCE [GENOMIC DNA].
     MEDLINE=94265501; PubMed=8205838;
RX
RA
     Delius H., Hofmann B.;
     "Primer-directed sequencing of human papillomavirus types.";
RT
```

RESULT 1

```
RL
    Curr. Top. Microbiol. Immunol. 186:13-31(1994).
CC
    -!- FUNCTION: Transcriptional transactivator. Binds double stranded
CC
        DNA (By similarity).
CC
    -!- SUBCELLULAR LOCATION: Nuclear matrix-associated (By similarity).
CC
    -!- SIMILARITY: Belongs to the papillomaviruses E6 protein family.
CC
    -----
CC
    Copyrighted by the UniProt Consortium, see http://www.uniprot.org/terms
CC
    Distributed under the Creative Commons Attribution-NoDerivs License
CC
    ------
DR EMBL; X74466; CAA52496.1; -; Genomic DNA.
DR
    PIR; S36538; S36538.
    InterPro; IPR001334; E6.
DR
DR
    Pfam; PF00518; E6; 1.
    Activator; DNA-binding; Early protein; Metal-binding; Nuclear protein;
KW
KW
    Transcription; Transcription regulation; Zinc; Zinc-finger.
FT
    CHAIN
                1
                     157
                             Protein E6.
FT
                             /FTId=PRO 0000133332.
FT
    ZN FING
                    75
              39
                             Potential.
              112
FT
    ZN FING
                     148
                            Potential.
    SEQUENCE 157 AA; 17984 MW; E9EC735537733FDC CRC64;
SQ
                      52.5%; Score 53; DB 1; Length 157;
 Query Match
 Best Local Similarity 53.3%; Pred. No. 14;
          8; Conservative 1; Mismatches 6; Indels 0; Gaps
 Matches
                                                                   0;
Qу
          1 WEAAAREACCRECCA 15
            1: |||||||
         63 WKGHFVTACCRSCCA 77
Db
Search completed: June 19, 2006, 17:39:02
```

Job time : 243 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:32:24 ; Search time 24 Seconds

(without alignments)

68.154 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : PIR 80:*

1: pir1:* 2: pir2:* 3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		6				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1 2 3 4 5 6 7 8 9	53 49 47.5 47 46 46 46 46 46	52.5 48.5 47.0 46.5 45.5 45.5 45.5 45.5	157 115 676 156 166 191 465 498 571	2 2 2 1 2 2 2 2 2 2	S36538 A36113 G84663 W6WL47 S36485 I46412 S05311 A48203 S69210 T34516	E6 protein - human antileukoproteinas hypothetical prote E6 protein - human E6 protein - human keratin KAP5.4 - s indoleacetamide hy interleukin-14 pre protein kinase cak hypothetical prote
11	45	44.6	61	2	E82580	hypothetical prote
12 13	45 45	44.6 44.6	155 157	1 1	W6WL8 W6WL5	E6 protein - human E6 protein - human
						2

						76
14	45	44.6	157	1	W6WLB5	E6 protein - human
15	45	44.6	273	2	A43862	29K peripheral mem
16	45	44.6	369	2	G75460	hypothetical prote
17	44	43.6	161	2	536491	E6 protein - human
18	44	43.6	186	2	A45910	ultra-high-sulfur
19	44	43.6	188	2	JC6547	high sulfur protei
20	44	43.6	204	2	T08072	proteinase inhibit
21	44	43.6	251	2	AH3413	nitrogen fixation
22	44	43.6	254	2	B84901	hypothetical prote
23	44	43.6	299	2	C97102	hypothetical prote
24	44	43.6	370	1	S57347	Ca2+/calmodulin-de
25	44	43.6	374	1	S50193	Ca2+/calmodulin-de
26	44	43.6	496	2	F75257	hypothetical prote
27	44	43.6	994	2	A48849	Ca2+-transporting
28	44	43.6	1001	1	PWRBFC	Ca2+-transporting
29	44	43.6	1121	2	S30862	DNA dependent ATPa
30	43.5	43.1	126	2	I46489	cysteine-rich hair
31	43	42.6	169	1	S18946	ultra high-sulfur
32	43	42.6	217	2	T33353	hypothetical prote
33	43	42.6	221	2	C34768	ORF2 protein - Orf
34	43	42.6	233	2	S67947	alkyl hydroperoxid
35	43	42.6	399	2	B24698	formate dehydrogen
36	43	42.6	689	2	T08988	cadmium-transporti
37	43	42.6	711	2	A85352	cadmium-transporti
38	43	42.6	976	2	D96714	DNA-directed RNA p
39	42.5	42.1	931	2	H96527	protein F27J15.16
40	42	41.6	122	2	JC6548	high sulfur protei
41	42	41.6	223	2	B38346	ultra-high-sulfur
42	42	41.6	230	2	A38346	ultra-high-sulfur
43	42	41.6	247	2	T17311	hypothetical prote
44	42	41.6	327	2	C86452	protein F6N18.11 [
45	42	41.6	1212	2	B82809	exodeoxyribonuclea
46	42	41.6	2037	2	T16881	hypothetical prote
47	41	40.6	67	2	T37199	hypothetical prote
48	41	40.6	151	2	S60314	hair keratin cyste
49	41	40.6	164	2	T24272	hypothetical prote
50	41	40.6	169	2	T06062	hypothetical prote
	* **		-07	_	10002	mypochecical proce

```
RESULT 1
S36538
E6 protein - human papillomavirus type 12
C; Species: human papillomavirus type 12
C;Date: 20-Feb-1995 #sequence_revision 20-Feb-1995 #text_change 09-Jul-2004
C; Accession: S36538
R; Delius, H.; Hofmann, B.
submitted to the EMBL Data Library, August 1993
A; Description: Primer-directed sequencing of human papillomavirus types.
A; Reference number: S36469
A; Accession: S36538
A; Molecule type: DNA
A; Residues: 1-157 < DEL>
A; Cross-references: UNIPROT: P36803; UNIPARC: UPI00001383B8; EMBL: X74466;
NID:g396910; PIDN:CAA52496.1; PID:g396911
C; Superfamily: papillomavirus E6 protein
```

C; Keywords: DNA binding; early protein; nucleus; zinc finger

Query Match 52.5%; Score 53; DB 2; Length 157;

Best Local Similarity 53.3%; Pred. No. 2.9;
Matches 8; Conservative 1; Mismatches 6; Indels 0; Gaps 0;

Qу 1 WEAAAREACCRECCA 15 |: |||||| Db 63 WKGHFVTACCRSCCA 77

Search completed: June 19, 2006, 17:39:33

Job time : 40 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:56:29; Search time 13 Seconds

(without alignments)

29.497 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 96747 seqs, 22556637 residues

Total number of hits satisfying chosen parameters: 96747

Minimum DB seq length: 0

Maximum DB seg length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : Published Applications AA New:*

1: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US09_NEW_PUB.pep:*

2: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US06_NEW_PUB.pep:*

3: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US07_NEW_PUB.pep:*

4: /EMC Celerra SIDS3/ptodata/1/pubpaa/US08 NEW PUB.pep:*

5: /EMC Celerra SIDS3/ptodata/1/pubpaa/PCT NEW PUB.pep:*

6: /EMC Celerra SIDS3/ptodata/1/pubpaa/US10 NEW PUB.pep:*

7: /EMC Celerra SIDS3/ptodata/1/pubpaa/US11_NEW_PUB.pep:*

8: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US60 NEW PUB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			•				
Res	sult		% Query				
	No.	Score	Match	Length	DB	ID	Description
	1	48	47.5	4440	6	US-10-196-749-525	Sequence 525, App
	2	45.5	45.0	139	6	US-10-449-902-42349	Sequence 42349, A
	3	45	44.6	1129	6	US-10-527-411-42	Sequence 42, Appl
	4	45	44.6	1129	6	US-10-527-411-48	Sequence 48, Appl
	5	45	44.6	1129	6	US-10-527-411-52	Sequence 52, Appl
	6	45	44.6	1129	6	US-10-527-411-56	Sequence 56, Appl
	7	45	44.6	1132	6	US-10-527-411-46	Sequence 46, Appl
	8	45	44.6	1894	6	US-10-196-749-97	Sequence 97, Appl
	9	44	43.6	105	6	US-10-953-349-39499	Sequence 39499, A

```
10
                              US-10-953-349-8402
        44
             43.6
                      449
                           6
                                                           Sequence 8402, Ap
11
        44
             43.6
                      449
                              US-10-953-349-9264
                                                           Sequence 9264, Ap
12
        44
             43.6
                      483
                           6
                              US-10-953-349-8401
                                                           Sequence 8401, Ap
13
             43.6
        44
                      483
                           6
                              US-10-953-349-9263
                                                           Sequence 9263, Ap
14
        44
             43.6
                      485
                           6
                              US-10-953-349-8400
                                                           Sequence 8400, Ap
15
             43.6
        44
                      485
                              US-10-953-349-9262
                                                           Sequence 9262, Ap
16
             43.6
                      804
        44
                           7
                              US-11-293-697-4161
                                                           Sequence 4161, Ap
17
             43.6
                     1435
        44
                           6
                              US-10-196-749-581
                                                           Sequence 581, App
18
             43.6
                     1743
        44
                           6
                              US-10-196-749-451
                                                           Sequence 451, App
19
             42.6
                       21 7
        43
                              US-11-144-322-3
                                                           Sequence 3, Appli
20
                              US-10-449-902-55514
        43
             42.6
                      198 6
                                                           Sequence 55514, A
21
        43
             42.6
                      257
                              US-10-953-349-31818
                                                           Sequence 31818, A
22
        42
             41.6
                       29
                           1
                              US-09-949-925-229
                                                           Sequence 229, App
23
                      113
        42
             41.6
                           6
                              US-10-953-349-33908
                                                           Sequence 33908, A
24
        42
             41.6
                      113
                              US-10-953-349-37356
                                                           Sequence 37356, A
25
        42
             41.6
                      145
                           6
                              US-10-953-349-33907
                                                           Sequence 33907, A
26
        42
             41.6
                      152
                           6
                              US-10-953-349-37355
                                                           Sequence 37355, A
27
        42
             41.6
                      161
                           1
                              US-09-949-925-226
                                                           Sequence 226, App
28
        42
             41.6
                      217
                           6
                              US-10-449-902-39327
                                                           Sequence 39327, A
29
        42
             41.6
                      414
                           6
                              US-10-449-902-32815
                                                           Sequence 32815, A
30
        42
             41.6
                      414
                              US-10-449-902-37283
                                                           Sequence 37283, A
31
        42
             41.6
                      414
                              US-10-449-902-46357
                                                           Sequence 46357, A
32
      41.5
             41.1
                      436 6
                              US-10-449-902-37829
                                                           Sequence 37829, A
33
             40.6
                       60 6
                                                           Sequence 38433, A
        41
                              US-10-449-902-38433
34
        41
             40.6
                      167
                                                           Sequence 34493, A
                              US-10-953-349-34493
35
        41
             40.6
                      373
                           6
                              US-10-449-902-38114
                                                           Sequence 38114, A
36
        41
             40.6
                      373
                           6
                              US-10-449-902-47991
                                                           Sequence 47991, A
37
        41
             40.6
                      373
                              US-10-449-902-50488
                           6
                                                           Sequence 50488, A
38
        41
             40.6
                      429
                           6
                              US-10-953-349-34644
                                                           Sequence 34644, A
39
        41
             40.6
                      429
                           6
                              US-10-953-349-35589
                                                           Sequence 35589, A
40
        41
             40.6
                      553
                           6
                              US-10-953-349-34643
                                                           Sequence 34643, A
41
        41
             40.6
                      553
                           6
                              US-10-953-349-35588
                                                           Sequence 35588, A
42
         41
              40.6
                       599
                           6 US-10-953-349-34642
                                                            Sequence 34642, A
43
        41
             40.6
                      601
                           6
                              US-10-953-349-35587
                                                           Sequence 35587, A
44
        41
             40.6
                      643
                           7
                              US-11-251-673-5
                                                           Sequence 5, Appli
45
        41
             40.6
                      643
                              US-11-293-697-3832
                                                           Sequence 3832, Ap
46
        41
             40.6
                      685
                           7
                              US-11-293-697-3546
                                                           Sequence 3546, Ap
47
        41
             40.6
                      720
                           6
                              US-10-196-749-170
                                                           Sequence 170, App
                              US-11-101-316-38
48
        41
             40.6
                      720
                           7
                                                           Sequence 38, Appl
                              US-10-196-749-269
49
        41
             40.6
                     1300
                           6
                                                           Sequence 269, App
50
      40.5
             40.1
                     1066
                           6
                              US-10-511-455-2
                                                           Sequence 2, Appli
```

```
RESULT 1
US-10-196-749-525
; Sequence 525, Application US/10196749
; Publication No. US20060094864A1
; GENERAL INFORMATION:
  APPLICANT: Baker, Kevin P.
  APPLICANT:
               Chen, Jian
  APPLICANT:
               Desnoyers, Luc
  APPLICANT:
               Goddard, Audrey
  APPLICANT:
               Godowski, Paul J.
  APPLICANT:
               Gurney, Austin L.
```

```
APPLICANT: Pan, James
  APPLICANT: Smith, Victoria
  APPLICANT: Watanabe, Colin K.
   APPLICANT: Wood, William I.
   APPLICANT:
              Zhang, Zemin
   TITLE OF INVENTION: SECRETED AND TRANSMEMBRANE POLYPEPTIDES AND NUCLEIC
  TITLE OF INVENTION: ACIDS ENCODING THE SAME
  FILE REFERENCE: P3430R1C340
  CURRENT APPLICATION NUMBER: US/10/196,749
  CURRENT FILING DATE: 2002-07-16
   PRIOR APPLICATION NUMBER: 10/052586
  PRIOR FILING DATE: 2002-01-15
  PRIOR APPLICATION NUMBER: 60/059263
  PRIOR FILING DATE: 1997-09-18
  PRIOR APPLICATION NUMBER: 60/059266
  PRIOR FILING DATE: 1997-09-18
  PRIOR APPLICATION NUMBER: 60/062250
  PRIOR FILING DATE: 1997-10-17
  PRIOR APPLICATION NUMBER: 60/063120
  PRIOR FILING DATE: 1997-10-24
  PRIOR APPLICATION NUMBER: 60/063121
  PRIOR FILING DATE: 1997-10-24
  PRIOR APPLICATION NUMBER: 60/063486
  PRIOR FILING DATE: 1997-10-21
  PRIOR APPLICATION NUMBER: 60/063540
; PRIOR FILING DATE: 1997-10-28
; PRIOR APPLICATION NUMBER: 60/063541
  PRIOR FILING DATE: 1997-10-28
; PRIOR APPLICATION NUMBER: 60/063544
  PRIOR FILING DATE: 1997-10-28
  Prior Application data removed - See File Wrapper or PALM.
  NUMBER OF SEQ ID NOS: 612
; SEQ ID NO 525
   LENGTH: 4440
   TYPE: PRT
   ORGANISM: Homo Sapien
US-10-196-749-525
  Query Match
                         47.5%; Score 48; DB 6; Length 4440;
 Best Local Similarity
                         60.0%; Pred. No. 56;
 Matches
           9; Conservative 0; Mismatches
                                               6; Indels 0; Gaps
                                                                           0;
QУ
           3 AAAREACCRECCARA 17
              Db
        3098 AAACTACCTTCCGGA 3112
```

Search completed: June 19, 2006, 18:01:11 Job time: 20 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:56:15; Search time 125.5 Seconds

(without alignments)

62.746 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

Sequence: 1 WEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2097797 seqs, 463214858 residues

Total number of hits satisfying chosen parameters: 2097797

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

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6: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US11_PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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101	100.0	17	3	US-09-813-197-4	Sequence 4, Appli
101	100.0	17	4	US-10-126-752-1	Sequence 1, Appli
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101	100.0	17	4	US-10-339-712-4	Sequence 4, Appli
101	100.0	17	5	US-10-719-523-4	Sequence 4, Appli
101	100.0	17	5	US-10-772-164-1	Sequence 1, Appli
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- ; Patent No. US20020132248A1
- ; GENERAL INFORMATION:
- ; APPLICANT: Rothschild, Kenneth J.
- ; APPLICANT: Gite, Sadanand
- ; APPLICANT: Olejnik, Jerzy
- ; TITLE OF INVENTION: N-Terminal and C-Terminal Markers in Nascent Proteins
- ; FILE REFERENCE: AMBER-06819
- ; CURRENT APPLICATION NUMBER: US/09/973,145
- ; CURRENT FILING DATE: 2001-10-09
- ; PRIOR APPLICATION NUMBER: 09/382,950
- ; PRIOR FILING DATE: 1999-08-25

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Search completed: June 19, 2006, 18:00:47

Job time : 140.5 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:39:41; Search time 36.5 Seconds

(without alignments)

40.768 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

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Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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4	101	100.0	17	2	US-09-382-736B-4	Sequence 4, Appli
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7	101	100.0	17	2	US-09-880-132-48	Sequence 48, Appl
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    APPLICANT:
                Tsien, Roger Y.
    APPLICANT:
                Griffin, B. Albert
;
    TITLE OF INVENTION: TARGET SEQUENCES FOR SYNTHETIC MOLECULES
;
    NUMBER OF SEQUENCES:
;
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Fish & Richardson P.C.
      STREET: 4225 Executive Square, Suite 1400
      CITY: La Jolla
      STATE: CA
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COUNTRY: USA
       ZIP: 92037
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       COMPUTER: IBM Compatible
       OPERATING SYSTEM: Windows 95
       SOFTWARE: FastSEQ for Windows Version 2.0b
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/955,206
       FILING DATE: 21-OCT-1997
    ATTORNEY/AGENT INFORMATION:
      NAME: Haile, Ph.D., Lisa A.
       REGISTRATION NUMBER: 38,347
       REFERENCE/DOCKET NUMBER: 07257/060001
     TELECOMMUNICATION INFORMATION:
       TELEPHONE: 619/678-5070
       TELEFAX: 619/678-5099
   INFORMATION FOR SEQ ID NO: 1:
     SEQUENCE CHARACTERISTICS:
       LENGTH: 17 amino acids
       TYPE: amino acid
       TOPOLOGY: linear
     MOLECULE TYPE: peptide
     FEATURE:
       OTHER INFORMATION: the N-terminus is acetylated and
       OTHER INFORMATION: the C-terminus is amidated
US-08-955-206-1
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           17; Conservative 0; Mismatches 0; Indels 0; Gaps
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             Db
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; Patent No. 6831160
; GENERAL INFORMATION:
; APPLICANT: THE REGENTS OF THE UNIVERSITY OF CALIFORNIA
; APPLICANT: VALE, Ronald ; APPLICANT: THORN, Kurt
; APPLICANT: COOKE, Roger
; APPLICANT: MATUSKA, Marija
; APPLICANT: NABER, Nariman
  TITLE OF INVENTION: METHOD OF AFFINITY PURIFYING PROTEINS USING MODIFIED BIS-
ARSENICAL
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 FILE REFERENCE: REGEN1500-1
; CURRENT APPLICATION NUMBER: US/09/502,664A
; CURRENT FILING DATE: 2000-02-11
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn version 3.0
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Job time : 46.5 secs

OM protein - protein search, using sw model

June 19, 2006, 17:24:19; Search time 210.5 Seconds Run on:

(without alignments)

36.925 Million cell updates/sec

Title: US-10-772-164-1

Perfect score: 101

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Total number of hits satisfying chosen parameters: 2589679

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Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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4	101	100.0	17	4	AAM48100	Aam48100 Fluoresce
5	101	100.0	17	8	ADO06947	Ado06947 FLASH-bin
6	101	100.0	17	9	ADZ76895	Adz76895 RNA-tag f
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11	87	86.1	19	5	AAU75749	Aau75749 FLAsH pep
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13	81	80.2	19	7	ABR84531	Abr84531 FLAsH pep
14	76	75.2	595	8	ADQ76865	Adq76865 Adenosine
15	61	60.4	22	3	AAY88739	Aay88739 Core poly
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RESULT 1
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ID AAY05336 standard; peptide; 17 AA.
XX
AC AAY05336;
XX
DT 29-JUN-1999 (first entry)
XX
DE Target sequence peptide, SEQ ID NO. 1.
XX
```

```
KW
     Biarsenical compound; alpha-helix peptide; polypeptide purification;
KW
     immunoassay; crosslinking agent.
XX
OS
     Synthetic.
XX
PN
     WO9921013-A1.
XX
PD
     29-APR-1999.
XX
PF
     21-OCT-1998;
                    98WO-US022363.
XX
PR
     21-OCT-1997;
                    97US-00955050.
PR
     21-OCT-1997;
                    97US-00955206.
PR
     21-OCT-1997;
                    97US-00955859.
XX
PA
     (REGC ) UNIV CALIFORNIA.
XX
PΙ
     Tsien RY, Griffin AB;
XX
DR
     WPI; 1999-288410/24.
XX
PT
     Biarsenical compounds that react specifically with cysteine residues.
XX
PS
     Claim 10; Page 41; 77pp; English.
XX
CC
     This sequence represents a target alpha-helix sequence for the
CC
     biarsenical compounds (BC) of the invention, which are able to react
CC
     specifically with cysteine residues in a target sequence to generate a
CC
     detectable signal. The BCs are used: (i) as labels that allow
CC
     identification of carrier molecules, e.g. in polypeptide purification,
CC
     immunoassays or other chemical or biological assays, including labelling
CC
     in vivo, e.g. to identify, locate or quantify polypeptides or nucleic
CC
     acids); (ii) for attaching a polypeptide to a solid substrate; or (iii)
CC
     to induce a polypeptide domain to adopt a more nearly alpha-helical form,
CC
     e.g. a conformation that can bind a drug. Tetra-arsenical compounds
CC
     derived from the BCs are used to crosslink two binding partners, e.g. to
CC
     study the effect of dimerisation on signal transduction. The BCs react
CC
     specifically with Cys-containing targets, and can be engineered to have
     particular properties, especially ability to cross a biological membrane
CC
CC
     and absence of any self-fluorescence. Both the BC and its target sequence
     are small, and BC binding between them is reversible, e.g. by treatment
CC
CC
     with a dithiol. Particularly, the BC becomes fluorescent when bound to
CC
     its target, but with a significant red-shift from the fluorescence of
CC
     fluorescein, allowing detection with very low background
XX
SO
     Sequence 17 AA;
  Query Match
                          100.0%; Score 101; DB 2; Length 17;
  Best Local Similarity
                          100.0%; Pred. No. 1.8e-05;
  Matches
          17; Conservative
                               0; Mismatches
                                                 0;
                                                      Indels
                                                                 0; Gaps
                                                                             0;
Qу
            1 WEAAAREACCRECCARA 17
              1 WEAAAREACCRECCARA 17
Search completed: June 19, 2006, 17:31:56
Job time: 238.5 secs
```

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:24:38 ; Search time 203 Seconds

(without alignments)

77.464 Million cell updates/sec

Title: US-10-772-164-4

Perfect score: 94

Sequence: 1 AEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2849598 seqs, 925015592 residues

Total number of hits satisfying chosen parameters: 2849598

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : UniProt 7.2:*

1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		ક				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
				-		
1	51	54.3	278	2	Q6IGH0_DROME	Q6igh0 drosophila
2	51	54.3	502	2	Q9BGM9_9MAMM	Q9bgm9 tachyglossu
3	51	54.3	1408	2	Q381X2_9TRYP	Q381x2 trypanosoma
4	50	53.2	491	2	Q4T2B4_TETNG	Q4t2b4 tetraodon n
5	50	53.2	589	2	Q2JB13_9ACTO	Q2jb13 frankia sp.
6	49.5	52.7	602	2	Q75NZ5_CHLRE	Q75nz5 chlamydomon
7	49	52.1	62	2	Q4PN38_IXOSC	Q4pn38 ixodes scap
8	49	52.1	157	1	VE6_HPV12	P36803 human papil
9	49	52.1	1067	2	Q4QFE4_LEIMA	Q4qfe4 leishmania
10	48	51.1	358	2	Q3F8C0_9BURK	Q3f8c0 burkholderi
11	48	51.1	358	2	Q4BRZ7_BURVI	Q4brz7 burkholderi
12	47	50.0	168	1	VE6_HPV21	P28832 human papil
13	47	50.0	193	1	KR415_HUMAN	Q9byq5 homo sapien
14	47	50.0	199	2	Q3W2K8_9ACTO	Q3w2k8 frankia sp.
15	47	50.0	210	1	KRA47_HUMAN	Q9byr0 homo sapien
						-

```
16
           47
                            330 2 Q3E2M8 CHLAU
                 50.0
                                                                        03e2m8 chloroflexu
17
           47
                 50.0
                            399 2 Q3INU5 NATPD
                                                                        O3inu5 natronomona
                                                                  Q31nu5 natronomona
Q341e3 rhodopseudo
Q5rgi5 brachydanio
Q9vxg1 drosophila
P16343 lonchocarpu
Q52509 pseudomonas
Q4ckp2 trypanosoma
Q618g9 homo sapien
Q8mz55 drosophila
Q9byr2 homo sapien
Q28583 ovis aries
Q3vgi5 sphingopyxi
Q85299 orf virus.
Q2i0e2 oryza sativ
P91666 drosophila
Q6n8x8 rhodopseudo
P19922 bradyrhizob
Q4s3z6 tetraodon n
Q6pfs4 brachydanio
Q4t6w6 tetraodon n
Q4tbg6 tetraodon n
Q5upc9 mimivirus.
Q76ya2 aeromonas p
Q5ts12 anopheles g
Q9pxb1 human papil
Q40617 human papil
Q407p3 mus musculu
Q49146 capsicum an
Q3tdh8 mus musculu
                                                                       Q341e3 rhodopseudo
Q5rgi5 brachydanio
Q9vxg1 drosophila
P16343 lonchocarpu
                           438 2 Q341E3_RHOPA
18
          47
                 50.0
                          878 2
1959 1
19
                                      Q5RGI5_BRARE
          47
                 50.0
20
          47
                 50.0
                                       HANG DROME
21
          46
                 48.9
                         80 1
                                       IBB4 LONCA
22
                                       Q52509 PSESX
          46
                 48.9
                            88 2
23
                            95 2
                                       O4CKP2 TRYCR
           46
                 48.9
24
                                      KRA56 HUMAN
          46
                 48.9 129 1
                           161 2
                                       Q8MZ55 DROME
25
          46
                 48.9
                           186 1
191 2
26
                                       KRA45 HUMAN
          46
                 48.9
27
                                       Q28583 SHEEP
          46
                 48.9
28
          46
                 48.9
                            203 2
                                       Q3VGI5_9SPHN
29
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                 48.9
                            221 2
                                       Q85299 9POXV
30
          46
                 48.9
                            232 2
                                       Q2I0E2 ORYSA
                         412 2 P91666 DROME
31
          46
                 48.9
                           441 2
465 1
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32
          46
                 48.9
33
          46
                 48.9
                                       HYIN2 BRAJA
                                       Q4S3Z6 TETNG
34
          46
                 48.9
                            533 2
35
          46
                 48.9
                           757 2 Q6PFS4 BRARE
36
          46
                 48.9 1033 2 Q4T6W6 TETNG
37
          46
                 48.9 1063 2 Q4TBG6 TETNG
38
          45
                 47.9
                           100 1 YL053_MIMIV
                           117 2
140 2
39
          45
                47.9
                                       Q76YA2 9CAUD
40
          45
                47.9
                                      Q5TS12 ANOGA
41
          45
                47.9
                            155 2
                                      Q9PXB1 HPV08
42
          45
                47.9
                            157 2
                                       O40617 HPVR7
43
          45
                47.9
                            165 2
                                       Q9D7P3 MOUSE
44
          45
                 47.9
                            204 1
                                      IP22 CAPAN
                            216 2
45
          45
                                       Q3TDH8 MOUSE
                                                                       Q3tdh8 mus musculu
Q7rzm5 neurospora
                 47.9
                           233 2
250 2
46
          45
                 47.9
                                       Q7RZM5 NEUCR
47
          45
                 47.9
                                                                       Q3wib2 frankia sp.
                                      Q3WIB2_9ACTO
48
          45
                 47.9
                            262 2 Q4U5Z5_CAPAN
                                                                       Q4u5z5 capsicum an
49
                 47.9
          45
                            262 2 Q4ZIQ3 CAPAN
                                                                        Q4ziq3 capsicum an
50
          45
                 47.9
                            262 2 Q4ZIQ4 CAPAN
                                                                         Q4ziq4 capsicum an
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RESULT 1
Q6IGHO DROME
     O6IGHO DROME
                    PRELIMINARY;
                                   PRT;
                                           278 AA.
AC
     O6IGH0;
DT
     05-JUL-2004, integrated into UniProtKB/TrEMBL.
DT
     05-JUL-2004, sequence version 1.
DT
     07-FEB-2006, entry version 8.
DE
     HDC06306.
GN
     ORFNames=HDC06306;
OS
     Drosophila melanogaster (Fruit fly).
OC
     Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
     Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
OC
     Ephydroidea; Drosophilidae; Drosophila.
OX
     NCBI_TaxID=7227;
RN
     [1]
RP
    NUCLEOTIDE SEQUENCE.
     PubMed=14709175; DOI=10.1186/gb-2003-5-1-r3;
RX
     Hild M., Beckmann B., Haas S.A., Koch B., Solovyev V., Busold C.,
RA
```

```
Fellenberg K., Boutros M., Vingron M., Sauer F., Hoheisel J.D.,
RA
    Paro R.;
RT
    "An integrated gene annotation and transcriptional profiling approach
RT
    towards the full gene content of the Drosophila genome.";
    Genome Biol. 5:RESEARCH0003.1-RESEARCH0003.17(2003).
RL
CC
    -!- MISCELLANEOUS: The sequence shown here is derived from an
CC
       EMBL/GenBank/DDBJ third party annotation (TPA) entry.
CC
    Copyrighted by the UniProt Consortium, see http://www.uniprot.org/terms
CC
CC
    Distributed under the Creative Commons Attribution-NoDerivs License
CC
    ______
    EMBL; BK003796; DAA02494.1; -; Genomic_DNA.
DR
DR
    InterPro; IPR013032; EGF_like_reg.
DR
    PROSITE; PS00022; EGF 1; UNKNOWN 1.
SO
    SEQUENCE 278 AA; 32016 MW; 06E7253102FE5BF1 CRC64;
 Query Match
                      54.3%; Score 51; DB 2; Length 278;
 Best Local Similarity 87.5%; Pred. No. 44;
 Matches
          7; Conservative 0; Mismatches
                                           1; Indels 0; Gaps
                                                                  0;
          9 CCRECCAR 16
Qу
            111111
Db
        250 CCRECCCR 257
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Search completed: June 19, 2006, 17:39:13

Job time : 214 secs

RA

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:32:24 ; Search time 24 Seconds

(without alignments)

68.154 Million cell updates/sec

Title: US-10-772-164-4

Perfect score: 94

Sequence: 1 AEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : PIR 80:*

1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		ř				
Result		Query				
No.	Score	Match	Length	DB 	ID	Description
1	49	52.1	157	2	S36538	E6 protein - human
2	46	48.9	191	2	146412	keratin KAP5.4 - s
3	46	48.9	221	2	C34768	ORF2 protein - Orf
4	46	48.9	273	2	A43862	29K peripheral mem
5	46	48.9	465	2	S05311	indoleacetamide hy
6	46	48.9	571	2	S69210	protein kinase cak
7	45	47.9	204	2	T08072	proteinase inhibit
8	45	47.9	370	1	S57347	Ca2+/calmodulin-de
9	45	47.9	374	1	S50193	Ca2+/calmodulin-de
10	44	46.8	188	2	JC6547	high sulfur protei
11	44	46.8	217	2	T33353	hypothetical prote
12	44	46.8	251	2	AH3413	nitrogen fixation
13	44	46.8	254	2	B84901	hypothetical prote

14	44	46.8	496	2	F75257	hypothetical prote
15	44	46.8	689	2	T08988	cadmium-transporti
16	44	46.8	711	2	A85352	cadmium-transporti
17	44	46.8	994	2	A48849	Ca2+-transporting
18	44	46.8	1001	1	PWRBFC	Ca2+-transporting
19	43.5	46.3	126	2	146489	cysteine-rich hair
20	43.5	46.3	229	2	S60454	glucose starvation
21	43	45.7	26	2	C39414	electron transport
22	43	45.7	156	1	W6WL47	E6 protein - human
23	43	45.7	157	1	W6WL5	E6 protein - human
24	43	45.7	157	1	W6WLB5	E6 protein - human
25	43	45.7	169	1	S18946	ultra high-sulfur
26	43	45.7	186	2	A45910	ultra-high-sulfur
27	43	45.7	233	2	S67947	alkyl hydroperoxid
28	43	45.7	399	2	B24698	formate dehydrogen
29	42.5	45.2	101	2	JQ0877	cyc02 protein prec
30	42	44.7	122	2	JC6548	high sulfur protei
31	42	44.7	166	2	S36485	E6 protein - human
32	42	44.7	223	2	B38346	ultra-high-sulfur
33	42	44.7	230	2	A38346	ultra-high-sulfur
34	42	44.7	327	2	C86452	protein F6N18.11 [
35	42	44.7	619	2	C96714	unknown protein T6
36	42	44.7	860	2	A96717	unknown protein, 4
37	42	44.7	997	2	S33754	glutamate receptor
38	42	44.7	2037	2	T16881	hypothetical prote
39	41	43.6	67	2	T37199	hypothetical prote
40	41	43.6	151	2	S60314	hair keratin cyste
41	41	43.6	155	1	W6WL8	E6 protein - human
42	41	43.6	161	2	S36491	E6 protein - human
43	41	43.6	164	2	T24272	hypothetical prote
44	41	43.6	169	2	T06062	hypothetical prote
45	41	43.6	188	2	T15651	hypothetical prote
46	41	43.6	199	2	T48099	hypothetical prote
47	41	43.6	352	2	S11926	cellulose 1,4-beta
48	41	43.6	369	2	F69407	iron-sulfur cluste
49	41	43.6	452	2	G86170	hypothetical prote
50	41	43.6	508	2	T22836	hypothetical prote

E6 protein - human papillomavirus type 12 C;Species: human papillomavirus type 12 C;Date: 20-Feb-1995 #sequence_revision 20-Feb-1995 #text_change 09-Jul-2004 C;Accession: S36538 R;Delius, H.; Hofmann, B. submitted to the EMBL Data Library, August 1993 A;Description: Primer-directed sequencing of human papillomavirus types. A;Reference number: S36469 A;Accession: S36538 A;Molecule type: DNA A;Residues: 1-157

A; Cross-references: UNIPROT: P36803; UNIPARC: UPI00001383B8; EMBL: X74466; NID: q396910: PIDN: CAA52496 1: PID: q396911

NID:g396910; PIDN:CAA52496.1; PID:g396911 C; Superfamily: papillomavirus E6 protein

RESULT 1

C; Keywords: DNA binding; early protein; nucleus; zinc finger

Query Match 52.1%; Score 49; DB 2; Length 157;

Best Local Similarity 87.5%; Pred. No. 11;

Matches 7; Conservative 0; Mismatches 1; Indels 0; Gaps 0;

Qy 8 ACCRECCA 15 |||| ||| Db 70 ACCRSCCA 77

Search completed: June 19, 2006, 17:39:37

Job time : 28 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:56:29; Search time 13 Seconds

(without alignments)

29.497 Million cell updates/sec

Title: US-10-772-164-4

Perfect score: 94

Sequence: 1 AEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 96747 seqs, 22556637 residues

Total number of hits satisfying chosen parameters: 96747

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : Published Applications AA New:*

1: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US09 NEW PUB.pep:*

2: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US06_NEW_PUB.pep:*

3: /EMC Celerra SIDS3/ptodata/1/pubpaa/US07 NEW PUB.pep:*

4: /EMC Celerra SIDS3/ptodata/1/pubpaa/US08 NEW PUB.pep:*

5: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/PCT NEW PUB.pep:*

6: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US10_NEW_PUB.pep:*

7: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US11_NEW_PUB.pep:*

8: /EMC_Celerra_SIDS3/ptodata/1/pubpaa/US60_NEW_PUB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		ક				
Result		Query				
No.	Score	Match	Length	DB	ID	Description
1	49	52.1	1129	6	US-10-527-411-42	Sequence 42, Appl
2	49	52.1	1129	6	US-10-527-411-48	Sequence 48, Appl
3	49	52.1	1129	6	US-10-527-411-52	Sequence 52, Appl
4	49	52.1	1129	6	US-10-527-411-56	Sequence 56, Appl
5	49	52.1	1132	6	US-10-527-411-46	Sequence 46, Appl
6	48	51.1	1894	6	US-10-196-749-97	Sequence 97, Appl
7	48	51.1	4440	6	US-10-196-749-525	Sequence 525, App
8	47	50.0	1435	6	US-10-196-749-581	Sequence 581, App
9	46	48.9	1743	6	US-10-196-749-451	Sequence 451, App

10	44	46.8	257	6	US-10-953-349-31818	Sequence	31818, A
11	44	46.8	449	6	US-10-953-349-8402		8402, Ap
12	44	46.8	449	6	US-10-953-349-9264	Sequence	_
13	44	46.8	483	6	US-10-953-349-8401	Sequence	
14	44	46.8	483	6	US-10-953-349-9263	Sequence	
15	44	46.8	485	6	US-10-953-349-8400	Sequence	8400, Ap
16	44	46.8	485	6	US-10-953-349-9262		9262, Ap
17	43	45.7	21	7	US-11-144-322-3	Sequence	3, Appli
18	43	45.7	198	6	US-10-449-902-55514		55514, A
19	43	45.7	1300	6	US-10-196-749-269	Sequence	269, App
20	42	44.7	29	1	US-09-949-925-229		229, App
21	42	44.7	161	1	US-09 - 949-925-226	Sequence	226, App
22	42	44.7	217	6	US-10-449-902-39327		39327, A
23	42	44.7	1776	6	US-10-933-854-3	Sequence	3, Appli
24	41.5	44.1	113	6	US-10-953-349-33908	Sequence	33908, A
25	41.5	44.1	113	6	US-10-953-349-37356	Sequence	37356, A
26	41.5	44.1	145	6	US-10-953-349-33907	Sequence	33907, A
27	41.5	44.1	152	6	US-10-953-349-37355	Sequence	37355, A
28	41	43.6	167	6	US-10-953-349-34493	Sequence	34493, A
29	41	43.6	429	6	US-10 - 953-349-34644		34644, A
30	.41	43.6	429	6	US-10-953-349-35589		35589, A
31	41	43.6	553	6	US-10-953-349-34643		34643, A
32	41	43.6	553	6	US-10-953-349-35588	Sequence	35588, A
33	41	43.6	599	6	US-10-953-349-34642	Sequence	34642, A
34	41	43.6	601	6	US-10-953-349-35587	Sequence	35587, A
35	41	43.6	643	7	US-11-251-673-5	Sequence	5, Appli
36	41	43.6	643	7	US-11-293-697-3832	Sequence	3832, Ap
37	41	43.6	685	7	US-11-293-697-3546	Sequence	3546, Ap
38	41	43.6	720	6	US-10-196-749-170	Sequence	170, App
39	41	43.6	720	7	US-11-101-316-38	Sequence	38, Appl
40	40.5	43.1	1066	6	US-10-511-455-2		2, Appli
41	40	42.6	181	6	US-10-953-349-10362	Sequence	10362, A
42	40	42.6	183	6	US-10-449-902-30401	Sequence	30401, A
43	40	42.6	183	6	US-10-449-902-45055	Sequence	45055, A
44	40	42.6	183	6	US-10-449-902-51021	Sequence	51021, A
45	40	42.6	201	6	US-10-953-349-3609	Sequence	3609, Ap
46	40	42.6	227	6	US-10-449-902-39040	Sequence	39040, A
47	40	42.6	282	7	US-11-293-697-3671	Sequence	3671, Ap
48	40	42.6	306	6	US-10-953-349-3608	Sequence	3608, Ap
49	40	42.6	331	6	US-10-953-349-3607	Sequence	3607, Ap
50	40	42.6	520	6	US-10-449-902-43105		43105, A
					ALIGNMENTS		

RESULT 1

US-10-527-411-42

- ; Sequence 42, Application US/10527411
- ; Publication No. US20060110410A1
- ; GENERAL INFORMATION:
- ; APPLICANT: Shone, Clifford
- ; APPLICANT: Foster, Keith Alan ; APPLICANT: Chaddock, John
- ; APPLICANT: Marks, Philip
- ; APPLICANT: Sutton, J. Mark
- ; APPLICANT: Stancombe, Patrick
- ; APPLICANT: Wayne, Jonathan
- ; TITLE OF INVENTION: Recombinant Toxin Fragments

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; FILE REFERENCE: 1581.0130005
; CURRENT APPLICATION NUMBER: US/10/527,411
; CURRENT FILING DATE: 2005-03-11
; PRIOR APPLICATION NUMBER: PCT/GB2003/003824
; PRIOR FILING DATE: 2003-09-12
  PRIOR APPLICATION NUMBER: US 10/241,596
; PRIOR FILING DATE: 2002-09-12
; NUMBER OF SEQ ID NOS: 175
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO 42
   LENGTH: 1129
   TYPE: PRT
   ORGANISM: Clostridium botulinum
US-10-527-411-42
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                         52.1%; Score 49; DB 6; Length 1129;
  Best Local Similarity 58.8%; Pred. No. 14;
                             3; Mismatches
 Matches
          10; Conservative
                                               4; Indels
                                                            0; Gaps
                                                                          0;
           1 AEAAAREACCRECCARA 17
Qу
             |||||:|| :| |:|
Db
         872 AEAAAKEAAAKEAAAKA 888
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Search completed: June 19, 2006, 18:01:13

Job time : 15 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:56:15; Search time 125.5 Seconds

(without alignments)

62.746 Million cell updates/sec

Title: US-10-772-164-4

Perfect score: 94

Sequence: 1 AEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2097797 seqs, 463214858 residues

Total number of hits satisfying chosen parameters: 2097797

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : Published Applications AA Main:*

1: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US07_PUBCOMB.pep:*

2: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US08_PUBCOMB.pep:*

3: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US09_PUBCOMB.pep:*

4: /EMC Celerra SIDS3/ptodata/2/pubpaa/US10A_PUBCOMB.pep:*

5: /EMC_Celerra_SIDS3/ptodata/2/pubpaa/US10B_PUBCOMB.pep:*

6: /EMC_Celerra SIDS3/ptodata/2/pubpaa/US11 PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	Query Match	Length	DB	ID	Description
1	94	100.0	17	3	US-09-880-149-49	Sequence 49, Appl
2	94	100.0	17	3	US-09-880-132-49	Sequence 49, Appl
3	94	100.0	17	4	US-10-126-752-4	Sequence 4, Appli
4	94	100.0	17	4	US-10-345-281-49	Sequence 49, Appl
5	94	100.0	17	5	US-10-772-164-4	Sequence 4, Appli
6	90	95.7	17	3	US-09-973-145-3	Sequence 3, Appli
7	90	95.7	17	3	US-09-880-149-48	Sequence 48, Appl
8	90	95.7	17	3	US-09-880-132-48	Sequence 48, Appl
9	90	95.7	17	3	US-09-813-197-4	Sequence 4, Appli
10	90	95.7	17	4	US-10-126-752-1	Sequence 1, Appli
11	90	95.7	17	4	US-10-174-368A-3	Sequence 3, Appli

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- ; Sequence 49, Application US/09880149
- ; Patent No. US20020146843A1
- ; GENERAL INFORMATION:
- ; APPLICANT: Kenten, John
- ; APPLICANT: Roberts, Steven
- ; TITLE OF INVENTION: CONTROLLING PROTEIN LEVELS IN EUCARYOTIC ORGANISMS
- ; FILE REFERENCE: 2757-5
- ; CURRENT APPLICATION NUMBER: US/09/880,149
- ; CURRENT FILING DATE: 2001-06-14
- ; PRIOR APPLICATION NUMBER: 09/406,781
- PRIOR FILING DATE: 1999-09-28
- ; PRIOR APPLICATION NUMBER: 60/119,851

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; PRIOR FILING DATE: 1999-02-12
; NUMBER OF SEQ ID NOS: 67
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 49
   LENGTH: 17
   TYPE: PRT
   ORGANISM: Unknown Organism
   FEATURE:
   OTHER INFORMATION: Description of Unknown Organism: example peptide
US-09-880-149-49
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                        100.0%; Score 94; DB 3; Length 17;
 Best Local Similarity 100.0%; Pred. No. 9.3e-05;
 Matches 17; Conservative 0; Mismatches 0; Indels
                                                              0; Gaps
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Qу
           1 AEAAAREACCRECCARA 17
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Db
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Search completed: June 19, 2006, 18:00:54

Job time : 132.5 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:39:41; Search time 36.5 Seconds

(without alignments)

40.768 Million cell updates/sec

Title: US-10-772-164-4

Perfect score: 94

Sequence: 1 AEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 650591 seqs, 87530628 residues

Total number of hits satisfying chosen parameters: 650591

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : Issued_Patents AA:*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result		% Query				
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1	94	100.0	17	1	US-08-955-206-4	Sequence 4, Appli
2	94	100.0	17	2	US-08-955-050-4	Sequence 4, Appli
3	94	100.0	17	2	US-09-406-781-49	Sequence 49, Appl
4	94	100.0	17	2	US-09-372-338-4	Sequence 4, Appli
5	94	100.0	17	2	US-09-880-132-49	Sequence 49, Appl
6	94	100.0	17	2	US-10-126-752-4	Sequence 4, Appli
7	90	95.7	17	1	US-08-955-206-1	Sequence 1, Appli
8	90	95.7	17	2	US-08-955-050-1	Sequence 1, Appli
9	90	95.7	17	2	US-09-382-950-3	Sequence 3, Appli
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RESULT 1
US-08-955-206-4
; Sequence 4, Application US/08955206
; Patent No. 5932474
; GENERAL INFORMATION:
; APPLICANT: Tsien, Roger Y.
; APPLICANT: Griffin, B. Albert
; TITLE OF INVENTION: TARGET SEQUENCES FOR SYNTHETIC MOLECULES
; NUMBER OF SEQUENCES: 4
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Fish & Richardson P.C.
; STREET: 4225 Executive Square, Suite 1400
; CITY: La Jolla
```

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STATE: CA
       COUNTRY: USA
       ZIP: 92037
     COMPUTER READABLE FORM:
       MEDIUM TYPE: Diskette
       COMPUTER: IBM Compatible
      OPERATING SYSTEM: Windows 95
      SOFTWARE: FastSEO for Windows Version 2.0b
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/955,206
      FILING DATE: 21-OCT-1997
    ATTORNEY/AGENT INFORMATION:
      NAME: Haile, Ph.D., Lisa A.
      REGISTRATION NUMBER: 38,347
      REFERENCE/DOCKET NUMBER: 07257/060001
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 619/678-5070
       TELEFAX: 619/678-5099
   INFORMATION FOR SEO ID NO: 4:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 17 amino acids
      TYPE: amino acid
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
US-08-955-206-4
  Query Match
                         100.0%; Score 94; DB 1; Length 17;
  Best Local Similarity 100.0%; Pred. No. 6.3e-05;
  Matches 17; Conservative 0; Mismatches 0; Indels
                                                              0; Gaps
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Qу
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Search completed: June 19, 2006, 17:41:14 Job time : 39.5 secs

OM protein - protein search, using sw model

Run on: June 19, 2006, 17:24:19; Search time 210.5 Seconds

(without alignments)

36.925 Million cell updates/sec

Title: US-10-772-164-4

Perfect score: 94

Sequence: 1 AEAAAREACCRECCARA 17

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2589679 seqs, 457216429 residues

Total number of hits satisfying chosen parameters: 2589679

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 1000 summaries

Database : A Geneseq 8:*

1: geneseqp1980s:*

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9: geneseqp2005s:*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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DT
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                  (first entry)
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KW
     Biarsenical compound; alpha-helix peptide; polypeptide purification;
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PR
     21-OCT-1997;
                    97US-00955206.
PR
     21-OCT-1997;
                    97US-00955859.
XX
PA
     (REGC ) UNIV CALIFORNIA.
XX
PΙ
     Tsien RY, Griffin AB;
XX
DR
     WPI; 1999-288410/24.
XX
PT
     Biarsenical compounds that react specifically with cysteine residues.
XX
PS
     Claim 10; Page 42; 77pp; English.
XX
CC
     This sequence represents a target alpha-helix sequence for the
CC
     biarsenical compounds (BC) of the invention, which are able to react
CC
     specifically with cysteine residues in a target sequence to generate a
CC
     detectable signal. The BCs are used: (i) as labels that allow
CC
     identification of carrier molecules, e.g. in polypeptide purification,
CC
     immunoassays or other chemical or biological assays, including labelling
CC
     in vivo, e.g. to identify, locate or quantify polypeptides or nucleic
CC
     acids); (ii) for attaching a polypeptide to a solid substrate; or (iii)
CC
     to induce a polypeptide domain to adopt a more nearly alpha-helical form,
CC
     e.g. a conformation that can bind a drug. Tetra-arsenical compounds
CC
     derived from the BCs are used to crosslink two binding partners, e.g. to
CC
     study the effect of dimerisation on signal transduction. The BCs react
CC
     specifically with Cys-containing targets, and can be engineered to have
CC
     particular properties, especially ability to cross a biological membrane
CC
     and absence of any self-fluorescence. Both the BC and its target sequence
     are small, and BC binding between them is reversible, e.g. by treatment
CC
CC
     with a dithiol. Particularly, the BC becomes fluorescent when bound to
CC
     its target, but with a significant red-shift from the fluorescence of
CC
     fluorescein, allowing detection with very low background
XX
SO
     Sequence 17 AA;
  Query Match
                          100.0%; Score 94; DB 2; Length 17;
  Best Local Similarity
                         100.0%; Pred. No. 0.00011;
 Matches
                                0; Mismatches 0; Indels
          17; Conservative
                                                                 0; Gaps
                                                                             0;
Qу
           1 AEAAAREACCRECCARA 17
              Db
           1 AEAAAREACCRECCARA 17
Search completed: June 19, 2006, 17:32:01
Job time: 215.5 secs
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